Contribution from the U. S. Army Quartermaster R & E Command.

Pioneering Research Division, Natick. Massace...

Synthesis of Organolead Arsonates and Arsinates

TECHNICAL LIBRARY

OUARTERMASTER RESEARCH &

ENGINEERING CENTER

Received March 14, 1962

BY MALCOLM C. HENRY

THE CONTROL OF THE PARTY NATICK, MASS.

A series of new organolead arsonates, =PbOAs(O)O-, and organolead arsinates, =PbOAs(O)=, has been prepared. Alternate methods of synthesis are discussed in connection with the relative ease of cleavage of leadcarbon and lead-halogen bonds. The properties of these compounds are discussed and a bactoness of ditto, foregoros permollo knodbiljishim obsorodusnost

Organometallic compounds containing M-O-M units (M = metal or metalloid atom) are well known for molecules where M atoms are identical and with M-O-M' units where M and M' are dissimilar but of the same group or subgroup in the periodic table. Less well known are compounds containing a group IVA element connected via oxygen to a group VA element other than nitrogen, phosphorus, or carbon.

The synthesis of M-O-As type compounds, for example, where M is silicon,1-8 germanium,4 or tin5-7 has been described only recently. The corresponding lead compounds have not been investigated for the case where lead is in its tetravalent state.

Synthesis of Organolead Arsinates.—The metathesis reaction between sodium dimethylarsinate and triphenyllead chloride was successful when effected in refluxing methanol. The same reaction when carried out in benzene resulted in yields of less than 1%, in accordance with the equation is at this equation are the large to be a subject to

 $(CH_3)_2As(O)ONa + (C_6H_5)_3PbCl \longrightarrow$ $(C_6H_5)_3PbOAs(O)(CH_3)_2(I) + NaCl$

The monosubstitution product (II) resulting from the reaction of sodium dimethylarsinate with diphenyllead dichloride was insoluble in dimethylformamide.

 $(CH_3)_2As(O)ONa + (C_6H_5)_2PbCl_2 \longrightarrow$

 $(C_6H_5)_2(Cl)PbOAs(O)(CH_3)_2(II) + NaCl$

The insolubility of II in dimethylformamide undoubtedly was reponsible for the fact that longer reaction times in the presence of excess sodium dimethylarsinate failed to produce the disubstituted product.

Ion Cleavage of Tetraphenyllead.—The electronegativity of the group IVA elements is known to vary widely. Relative values for carbon, 2.55; silicon, 1.76; germanium, 1.40; tin, 1.37; and lead, 1.138 are related to the observed differences in reactivities of compounds containing group IVA elements bonded directly to carbon. Large differences in reactivity have been noted even between tetraphenyltin and tetraphenyllead and the latter has been reported to be 60 times more sensitive to acid cleavage. 98,6

The reaction of tetraphenyllead with HCl is stepwise and can be controlled so as to isolate either the monohalide or the relatively insoluble dihalide. 10 It has not been possible to extend this substitution process to the tri- or tetrahalide. It has been suggested that because of the low dielectric constant of benzene these reactions may take place by a molecular attack of HX on tetraphenyllead.9a

Many organolead compounds have been prepared by the reaction of tetrasubstituted lead compounds with mineral or organic acids. 11 The salt-like character of these compounds is suggested by the fact that compounds derived from weak acids generally are more soluble in organic solvents than those formed from strong acids. This is complicated by the case of monosubstituted acid salts of tetraphenyllead because of the sensitivity to cleavage of the second phenyl group by acids to

⁽¹⁾ R. M. Kary and K. C. Prisch, J. Am. Chem. Soc., 79, 2140 (1957).

⁽²⁾ M. Schmidt and H. Schmidbaur, Angew. Chem., 71, 553 (1959).

⁽³⁾ B. L. Chamberland and A. G. MacDiarmid, J. Am. Chem. Soc., 82, 4542 (1960); 83, 549 (1961).

⁽⁴⁾ M. Schmidt, I. Rudisch, and H. Schmidbaur, Ber., 94, 2451 $(1961)_{-}$

⁽⁵⁾ A. W. Walde, H. E. Van Essen, and T. W. Zbornik, U. S. Pat. No. 2,762,821 (1956); Chem. Abstr., 51, 4424 (1957).

⁽⁶⁾ E. Rochow, D. Seyferth, and A. C. Smith, J. Am. Chem. Soc., 75, 3099 (1953).

⁽⁷⁾ B. L. Chamberland and A. G. MacDiarmid, J. Chem. Soc., 445 (1960).

⁽⁸⁾ R. K. Sheline and K. S. Pitzer, J. Chem. Phys., 18, 595 (1950).

^{(9) (}a) K. Jim-Young, Dissertation Abstr., 21, 3271 (1961); (b) R. E. Dessy and K. Jim-Young, J. Am. Chem. Soc., 83, 1167 (1961). (10) H. Gilman and J. D. Robinson, ibid., 51, 3112 (1929).

⁽¹¹⁾ R. W. Leeper, L. Summers, and H. Gilman, Chem. Rev., 54, 101 (1954):

yield the disubstituted acid salts. Triphenyllead chloride, for example, when allowed to react with glacial acetic acid, readily forms diphenylchlorolead acetate. Triphenyllead acetate when allowed to react with dry hydrochloric acid in benzene also forms diphenylchlorolead acetate initially; however, upon further reaction, the acetate ion is displaced to the almost complete insolubility of the latter in organic solvents.

Organolead Arsonates.—Two equivalents of benzylarsonic acid when allowed to react with triphenyllead chloride in methanol produced an organolead diarsonate, whereas if propylarsonic acid was used in equivalent molar quantities only the monosubstituted product was obtained, i.e.

$$(C_6H_5)_2PbCI + 2C_6H_5CH_2As(O)(OH)_2 \xrightarrow{CH_6OH}$$

$$(C_6H_5)_2Pb[OAs(O)(OH)CH_2C_6H_5]_2 (III) +$$

$$HCl + C_6H_6$$

or a first this file is the feature of this is

$$(C_6H_6)_8PbCl + CH_3CH_2CH_2As(O)(OH)_2 \longrightarrow$$
 $(C_6H_6)_6(Cl)PbOAs(O)(OH)CH_2CH_2CH_6(IV) + C_6H_6$

The use of pyridine as a solvent makes it possible to enhance the reaction by the formation of pyridine hydrochloride. Thus it was possible to synthesize the disubstituted organolead arsonates

$$(C_6H_6)_2PbCI + 2RAs(O)(OH)_2 \xrightarrow{C_6H_6N}$$

$$(C_6H_6)_2Pb[OAs(O)(OH)R]_2 +$$

$$C_6H_6 + C_5H_5N\cdot HCI$$

where $R = CH_2 = CHCH_2$ (V); $m-NO_2C_6H_4^-$ (VI); or $C_6H_5^-$ (VII). A similar reaction starting with tetraphenyllead instead of triphenyllead chloride using m-nitrophenylarsonic acid resulted also in the isolation of compound VI

$$(C_6H_5)_4P_5 + 2m \cdot NO_2C_6H_4As(O)(OH)_2 \xrightarrow{C_5H_5N} VI + 2C_6H_6$$

Reaction of triphenyllead chloride and methylarsonic acid in pyridine gave anomalous results. In this case two products, VIII and IX, were isolated, the former containing a ratio of lead to arsenic of 2:1. The second product was apparently the monosubstitution product which upon heating underwent dehydration to form the corresponding anhydride (X). Thus, it is conceivable that the following reaction occurred

$$(C_6H_5)_3PbCl + CH_3As(O)(OH)_2 \longrightarrow$$

$$[(C_6H_5)_2(Cl)PbO]_2As(O)CH_3 (VIII) +$$

$$(C_6H_5)_2(Cl)PbOAs(O)(OH)CH_3 (IX)$$

$$\begin{array}{c} -H_2O \\ IX \longrightarrow [(C_6H_6)_2(Cl)PbOAs(O)(CH_3)]_2O \end{array} (X)$$

Reaction between diphenyllead dichloride and disodium methylarsonate in dimethylformamide yields a white powdery material whose analysis indicates a lead—arsenic ratio of 1:1. It was not possible to characterize this compound further, although presumably the initial reaction involves a metathesis to form either a linear chain structure (XI)

$$(C_6H_5)_2PbCl_2 + CH_3As(O)(ONa)_2 \longrightarrow$$

$$\begin{bmatrix} O & C_6H_6 \\ -As-O-Pb-O & - \\ CH_3 & C_6H_5 \end{bmatrix} - (XI) + NaC$$

or a cyclic dimer (XII)

$$(C_6H_5)_2\mathrm{Pb}<(O-\mathrm{As}(O)(CH_3)-O)_2>\mathrm{Pb}(C_6H_5)_2\ (\mathrm{XII})$$

Reaction of Organolead Arsonates with Glacial Acetic Acid.—The reaction of organolead arsonates with glacial acetic acid makes it possible to replace the arsonate groups by acetate, yielding in these cases either diphenyllead diacetate or diphenylchlorolead acetate. This reaction quickly determines whether a mono- or disubstitution has taken place since diphenyllead diacetate crystallizes from glacial acetic acid upon standing and can be easily identified, whereas diphenylchlorolead acetate decomposes before melting and is insoluble in glacial acetic acid.

Since a minimum of two phenyl groups are assumed to be attached to the lead atom and since elemental analyses determine the lead-to-arsenic ratio, it is possible to identify all four groupings attached to the central lead atom. In the absence of this reaction, it is possible to have a Pb: As ratio of 1:1 and three possible configurations, either the monosubstituted $(C_6H_5)_2$ CIPbOAs-(O)=, the disubstituted cyclic product $(C_6H_5)_2$ Pb- $(OAs(O)(R)O]_2>Pb(C_6H_5)_2$, or a polymer. In its simplest sense this reaction with glacial acetic acid may be considered as a qualitative test for the presence of chloride.

These compounds, having no melting points up to 260°, all are insoluble in the common organic solvents, thus making purification difficult. As a result the elemental analyses are, more often than not, unacceptable by normal standards. It is felt, however, that they are sufficiently definitive to establish the identity of the compounds described. It was not possible to ascertain whether inter- or intramolecular dehydration was respon-

sible for the deviations noted. The infrared spectra of these compounds indicate that in certain cases this may be the case. Investigations currently are being carried out to clarify this point.

Infrared Studies.—Our studies have shown that the infrared spectra of organoarsonic acids have some useful absorptions which are pertinent to this Methyl-, phenyl-, benzyl-, β investigation. chlorovinyl-, allyl-, propyl-, m-nitrophenyl-, phydroxyphenyl-, and butylarsonic acids all show intense broad absorptions at ca. 2702 and 2381-2272 cm. -1 using potassium bromide pelleting techniques. It is assumed these absorptions are associated with the OH stretching vibrations, since these absorption bands are absent in the cases of triphenylarsine, arsenic trioxide, and methylarsine oxide.

Infrared absorption characteristic for aromatic compounds of lead appears as an intense narrow band at 1058 cm. -1.12 We have noted this band to be invariant for a large number of organolead compounds containing two to four phenyl groups. Other atoms attached to lead such as halogens or sulfur do not change the intensity of the absorption. During the course of this investigation, however, it was noted that with aryllead acetates and aryllead arsonates, that is, where there exists a Pb-O bond, the intensity of the 1058 cm.-1 absorption decreases markedly and in some cases almost disappears. Aryltin compounds have an analogous absorption at 1070 cm. -1 and it has been noted that in the case of aryltin arsonates this absorption also decreases and almost disappears.3

In general, the organolead arsonates show absorption bands in the 750-950 cm. -1 region which are absent in the starting materials. These bands presumably are associated with the Pb-O-As bonds. These bands are intense, sometimes multiple in nature, and broad. In the case of the organotin arsonates, new, intense, broad bands in the same general area of the spectrum are characteristic in appendix on these theorems A - 115 to other

The triphenyllead derivative of dimethylarsinic acid (compound I) shows two intense bands at 875 and 834 cm.-1, whereas the diphenylchlorolead derivative, compound II, shows an intense band at 790 cm.-1 that predominates the spectrum. Both compounds show the expected methyl and phenyl vibrations plus a weak phenyl-Pb absorption at 1058 cm - importation outlooks have

Table I summarizes the infrared spectra of the organolead arsonates (cm. -1). Column A lists the absorption bands formed which are present in the product but not in the reactants. Column B lists absorptions that have disappeared, but which were present in the parent arsonic acid.

i gradi gula Victoria. Lidafras las Caldras	TABLE I
(300,41).41	Organolead Parent arsonate arsonic acid (formed), (disappeared),
Com-	(formed), (disappeared),
pound	. Gem. to GIV. A. O., D., O. and cm. A. G Januari.
III 875	5, 855, 1015 (1986) 865, 2702, 2280 (1986) 40
VII 853	880,778 JEWE & COLOR
IV 3333 870	770, 915, 1220, 2702, 2280
	3,750, sabbe dika 768 min jepaka pot bazgi
)
	3, 840, 870, 880 \[\int 780, 940, 1215, 2702, 2280 \]
	5, 842, 870, 883 hay (a) desperant the singulative less
$\mathbf{X}_{\mathcal{H}}$ contains	ng by Am Out of Box beats ask Conforme Of
	0, 790 ng na sanggara 733, 820, 3350 na a pagaba

The formation of intense bands between 770 and 870 cm. -1 appears to be characteristic of Pb-O-As type compounds. Comparison with Sn-O-As spectra seems to indicate this latter structure absorbs between 800 and 900 cm. It should be realized that until more data are available these absorptions should not be interpreted as other than empirical characteristics of the products.

have been the Experimental and the second second

Melting points were taken with a Kofler hot stage. Organolead starting materials and reference materials tetraphenyllead,13 triphenyllead chloride and diphenyllead dichloride,10 triphenyllead iodide, triphenyllead acetate,14 and diphenyllead diacetate15 were prepared by published methods. The organosubstituted arsonic acids were generously supplied by American Smelting & Refining Co., Rahway, N. J., through the late R. Kary. Infrared spectra were obtained using a Perkin-Elmer Infracord spectro-

Triphenyllead Dimethylarsinate (I).—Sodium cacodylate trihydrate (Mallinckrodt) (10 mmoles) and triphenyllead iodide (10 mmoles) dissolved in 200 ml. of methanol were refluxed for 8 hr. The methanol was evaporated, the residue washed with water several times, and finally recrystallized twice from methanol; yield, pure triphenyllead dimethylarsinate, 4.2 g. (73%).

Anal. Calcd. for C20H21AsPbO2: C, 41,74; H, 3.65; Pb, 36.05; As, 13.04. Found: C, 41.58; H, 3.71; Pb, 35.20; As, 13.02.

The same reaction was carried out, as described above, in benzene; after 50 hr. reflux only 0.3 g. of triphenyllead dimethylarsinate was produced.

Triphenyllead dimethylarsinate refluxed together with excess glacial acetic acid produced diphenyllead diacetate

⁽¹²⁾ J. G. Noltes, M. C. Henry, and M. J. Janssen, Chem. Ind. (London), 298 (1959); bothery been harmin countrieved off

⁽¹³⁾ H. Gilman and J. Robinson, J. Am. Chem. Soc., 49, 2315 (1927).

⁽¹⁴⁾ R. Heap and B. C. Saunders, J. Chem. Soc., 658 (1951).

^{15 (15)} A. Polis, Ber., 20, 3331 (1887). A party domestic and the second of

quantitatively. The product was identified by mixed melting point with an authentic sample and comparison of infrared spectra of an authentic sample.

Diphenylchlorolead Dimethylarsinate (II).—Diphenyllead dichloride (10 mmoles), together with sodium cacodylate trihydrate (20 mmoles), was dissolved in 300 ml. of dimethylformamide and refluxed for 8 hr. A white insoluble precipitate formed on the sides of the flask during the reaction period. The precipitate was filtered, washed with several aliquots of water, and dried; yield, 4.1 g. (76.9%) of diphenylchlorolead dimethylarsinate.

Anal. Calcd for $C_{14}H_{16}O_2AsClPb$: C, 31.50; H, 3.00; Pb, 38.90; As, 14.03. Found: C, 30.83; H, 3.15; Pb, 39.00; As, 13.06.

An aliquot of diphenylchlorolead dimethylarsinate refluxed for several minutes with excess glacial acetic acid produced diphenylchlorolead acetate quantitatively.

Diphenyllead Bis-(benzylarsonate) (III).—Triphenyllead chloride (5 mmoles), together with benzylarsonic acid (10 mmoles), was dissolved in 400 ml. of methanol and refluxed for 16 hr. A white insoluble precipitate was produced. Filtration and washing with fresh solvent yielded 2.6 g. (35%) of diphenyllead bis-(benzylarsonate).

Anal. Calcd. for C₂₆H₂₆O₆As₂Pb: C, 39.50; H, 3.30; As, 18.90; Pb, 26.19. Found: C, 40.13; H, 3.81; As, 18.39; Pb, 25.9.

Reaction with glacial acetic acid produced diphenyllead

Diphenylchlorolead Propylarsonate (IV).—Triphenyllead chloride (5 mmoles), together with propylarsonic acid (5 mmoles), dissolved in 300 ml. of methanol and refluxed 12 hr. produced a white insoluble precipitate. Filtration and washing with successive portions of methanol and water yielded diphenylchlorolead propylarsonate, 2.5 g. (89% yield).

Anal. Calcd. for C₁₆H₁₈O₅AsClPb: C, 31.90; H, 3.20; As, 13.30; Pb, 36.80. Found: C, 31.40; H, 3.64; As, 13.37; Pb, 38.7.

Reaction with glacial acetic acid produced diphenylchlorolead acetate.

Diphenyllead Bis-(allylarsonate) (V).—Allylarsonic acid (10 mmoles), together with triphenyllead chloride (5 mmoles), was dissolved in 250 ml. of pyridine and heated at 70° for 24 hr. Cooling produced 2 g. of colorless crystals. Filtration and washing of the residue left 2.0 g. of diphenyllead bis-(allylarsonate), 35.4% yield.

Anal. Calcd. for C₁₈H₂₂O₆As₂Pb: C, 31.26; H, 3.19; As, 21.68; Pb, 30.00. Found: C, 30.91; H, 3.32; As, 21.72; Pb, 30.00.

Reaction with glacial acetic acid produced diphenyllead discrete

Diphenyllead Bis-(m-nitrophenylarsonate) (VI): m-Nitrophenylarsonic acid (10 mmoles), together with triphenyllead chloride (5 mmoles) in 300 ml. of pyridine, was heated at 70° for 24 hr. The pale yellow crystals that gradually formed during the reaction were filtered, washed with pyridine and methanol, and dried under vacuum; yield, 2.3 g. (53.8%) diphenyllead bis-(m-nitrophenyl arsonate).

Anal. Calcd. for $C_{24}H_{20}O_{10}As_2N_2Pb$: C, 33.76; H, 2.64; As, 17.54; Pb, 24.29. Found: C, 33.30; H, 2.65; As, 17.40; Pb, 23.85.

Reaction with glacial acetic acid produced diphenyllead

diacetate. (An alternate synthesis of VI was carried out using tetraphenyllead and *m*-nitrophenylarsonic acid with a similar stoichiometry of 1:2. The reaction was carried out in pyridine at 70° for 6 hr. and the product (VI) isolated as described above.)

Diphenyllead Bis-(phenylarsonate) (VII).—Phenylarsonic acid (10 mmoles), together with triphenyllead iodide (5 mmoles), dissolved in pyridine was heated 18 hr. at 80°. Upon standing overnight white crystals separated out of solution. Filtration and air drying yielded 1.5 g. (32.6%) of the dipyridine complex of diphenyllead bis-(phenylarsonate).

Anal. Calcd. for C₂₁H₃₂O₆As₂N₂Pb: C, 44.29; H, 3.58; As, 16.26; N, 3.03; Pb, 22.49. Found: C, 43.71; H, 3.49; As, 16.55; N, 2.91; Pb, 22.29.

An aliquot of the dipyridine complex boiled for I hr. with anhydrous methanol destroyed the complex to yield diphenyllead bis-(phenylarsonate).

Anal. Calcd for C₂₄H₂₂O₆As₂Pb: C, 37.74; H, 2.89; As, 19.63; Pb, 27.10. Found: C, 38.84; H, 3.38; As, 19.61; Pb, 26.1.

A second aliquot of the pyridine complex boiled for several minutes with glacial acetic acid yielded diphenyllead diacetate.

Anal. Calcd. for C₁₆H₁₆PbO₄: C, 40.08; H, 3.34; Pb, 43.25. Found: C, 40.08; H, 3.36; Pb, 43.20.

Reaction of Triphenyllead Chloride and Methylarsonic Acid.—Methylarsonic acid (0.02 mole) and triphenyllead chloride (0.02 mole) were dissolved in 200 ml. of pyridine and stirred for 12 hr. in an oil bath maintained at a temperature of 80-90°. The white crystalline plates that formed during the reaction were filtered and washed with chloroform; yield, 9.5 g. (88.7%) of diphenylchlorolead monobasic ester of methylarsonic acid (IX).

Anal. Calcd. for C₁₈H₁₄O₃AsClPb: C, 29.05; H, 2.61; As, 13.95; Pb, 38.6. Found: C, 28.74; H, 2.86; As, 13.62; Pb, 38.4.

Reaction with glacial acetic acid produced diphenylchloroacetate. Compound IX boiled in methanol for 30 min. slowly produced a precipitate no longer soluble in the solvent. This appears to be the corresponding anhydride compound X.

Anal. Calcd. for C₂₆H₂₆O₅As₂Cl₂Pb₂: C, 29.60; H, 2.47; As, 14.3; Pb, 39.3. Found: C, 29.59; H, 2.81; As, 14.05; Pb, 39.7.

Reaction with glacial acetic acid produced diphenylchlorolead acetate.

Evaporation of the original pyridine solution yielded a white powder which after washing with chloroform yielded 1.49 g. of a product which contains lead and arsenic in the ratio of 2:1. A compound such as compound VIII seems probable.

Anal. Calcd. for compound VIII (C₂₈H₂₈O₂AsCl₂Pb₂): C, 32.20; H, 2.47; As, 7.95; Pb, 44.5. Found: C, 30.32; H, 2.90; As, 7.88; Pb, 41.2.

Reaction with glacial acetic acid produced diphenylchlorolead acetate.

Reaction of Diphenyllead Dichloride and Disodium Methylarsonate.—Diphenyllead dichloride (0.01 mole) and disodium methylarsonate (0.01 mole) were dissolved in dimethylformamide and refluxed for 8 hr. During the reaction a white powdery precipitate gradually formed. The precipitate, filtered and washed with fresh solvent

921

several times, yielded $6.0\,\mathrm{g}$ of product insoluble in common organic solvents.

Anal. Calcd. for $(C_{13}H_{18}O_3AsPb)_x$ (XI or XII): C, 31.3; H, 2.61; As, 15.0; Pb, 41.5. Found: C, 28.48; H, 2.72; As, 13.87; Pb 41.8.

This product boiled together with glacial acetic acid yielded diphenyllead diacetate.

Acknowledgment.—The author is grateful to C. DiPietro of the Analytical Group of this Division for the elemental analyses.